

	Type	L #	Hits	Search Text	DBs	Time Stamp	Comments	Error Definition	Error Count
1	BRS	L1	22465	glycoprotein	USPAT; US-PGPUB; EPO; JPO; DERWENT	2002/03/24 15:08			0
2	BRS	L2	2397	(helicobacter adj pylori) or (H. adj pylori)	USPAT; US-PGPUB; EPO; JPO; DERWENT	2002/03/24 15:08			0
3	BRS	L3	233	urease same ((helicobacter adj pylori) or (H. adj pylori))	USPAT; US-PGPUB; EPO; JPO; DERWENT	2002/03/24 15:08			0
4	BRS	L4	4	glycoprotein same (urease same ((helicobacter adj pylori) or (H. adj pylori)))	USPAT; US-PGPUB; EPO; JPO; DERWENT	2002/03/24 15:08			0
5	BRS	L5	36	glycoprotein same whey same milk	USPAT; US-PGPUB; EPO; JPO; DERWENT	2002/03/24 15:11			0
6	BRS	L6	33	glycoprotein same albumin same egg	USPAT; US-PGPUB; EPO; JPO; DERWENT	2002/03/24 15:11			0
7	BRS	L7	2	((glycoprotein same whey same milk) or (glycoprotein same albumin same milk)) same (urease same ((helicobacter adj pylori) or (H. adj pylori)))	USPAT; US-PGPUB; EPO; JPO; DERWENT	2002/03/24 15:12			0
8	BRS	L8	3	glycoprotein same ((helicobacter adj pylori) or (H. adj pylori)) same inhibit\$ same colonization	USPAT; US-PGPUB; EPO; JPO; DERWENT	2002/03/24 15:13		Truncation overflow low.	1

	Type	L #	Hits	Search Text	DBs	Time Stamp	Comments	Error Definition	Error Count
9	BRS	L9	8	((helicobacter adj pylori) or (H. adj pylori)) same inhibitor same colonization	USPAT; US-PGPUB; EPO; JPO; DERWENT	2002/03/24 15:14			0

=> d his

(FILE 'HOME' ENTERED AT 15:21:03 ON 24 MAR 2002)

FILE 'MEDLINE, CAPLUS, BIOSIS, EMBASE, SCISEARCH, AGRICOLA'
ENTERED AT

15:21:34 ON 24 MAR 2002

L1 497584 S GLYCOPROTEIN
L2 77272 S HELICOBACTER PYLORI
L3 6694 S L2 (P) UREASE
L4 24 S L1 (P) L3
L5 9 DUPLICATE REMOVE L4 (15 DUPLICATES REMOVED)
L6 403 S L2 (P) COLONIZATION (P) INHIBIT?
L7 20 S L6 AND L1
L8 8 DUPLICATE REMOVE L7 (12 DUPLICATES REMOVED)
L9 6 S L8 NOT L5

=> log y

FILE 'HOME' ENTERED AT 15:21:03 ON 24 MAR 2002

=> file medline caplus biosis embase scisearch agricola

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ENTRY

SESSION

FULL ESTIMATED COST

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FILE 'MEDLINE' ENTERED AT 15:21:34 ON 24 MAR 2002

FILE 'CAPLUS' ENTERED AT 15:21:34 ON 24 MAR 2002

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FILE 'AGRICOLA' ENTERED AT 15:21:34 ON 24 MAR 2002

=> s glycoprotein

L1 497584 GLYCOPROTEIN

=> s helicobacter pylori

L2 77272 HELICOBACTER PYLORI

=> s l2 (p) urease

L3 6694 L2 (P) UREASE

=> s l1 (p) l3

L4 24 L1 (P) L3

=> duplicate remove l4

DUPLICATE PREFERENCE IS 'MEDLINE, CAPLUS, BIOSIS, EMBASE, SCISEARCH'

KEEP DUPLICATES FROM MORE THAN ONE FILE? Y/(N):n

PROCESSING COMPLETED FOR L4

L5 9 DUPLICATE REMOVE L4 (15 DUPLICATES REMOVED)

=> d l5 1-9 ibib abs

L5 ANSWER 1 OF 9 CAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER: 2001:760034 CAPLUS

DOCUMENT NUMBER: 135:278059

TITLE: Glycoprotein having inhibitory activity against
Helicobacter pylori colonization

INVENTOR(S): Kodama, Yoshikatsu; Kimura, Nobutake

PATENT ASSIGNEE(S): Ghen Corporation, Japan; Nisshin Flour Milling Co.,
Ltd.

SOURCE: Eur. Pat. Appl., 16 pp.

CODEN: EPXXDW

DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
EP 1145644	A2	20011017	EP 2001-400969	20010413
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO				
JP 2001294600	A2	20011023	JP 2000-113913	20000414
US 2001044120	A1	20011122	US 2001-833637	20010413
PRIORITY APPLN. INFO.:		JP 2000-113913	A	20000414
AB An inhibitor of ***Helicobacter*** ***pylori*** colonization in the stomach comprises as an active ingredient a ***glycoprotein*** which specifically binds to H. pylori ***urease***. This				

glycoprotein is isolated and purified from a ***glycoprotein***-contg. substance, esp. that derived from bovine milk whey or yolk of chicken eggs by affinity chromatog. using a column on which H. pylori ***urease*** is immobilized. The ***glycoprotein*** is able to effectively inhibit H. pylori colonization, and thus is useful for the prevention or treatment of diseases caused by infection of H. pylori such as peptic ulcers. A food and medicament comprising the inhibitor are also provided.

L5 ANSWER 2 OF 9 MEDLINE DUPLICATE 1
 ACCESSION NUMBER: 2000403971 MEDLINE
 DOCUMENT NUMBER: 20389972 PubMed ID: 10930371
 TITLE: Acid-dependent adherence of Helicobacter pylori urease to diverse polysaccharides.
 AUTHOR: Icatlo F C; Goshima H; Kimura N; Kodama Y
 CORPORATE SOURCE: Immunology Research Institute, Ghen Corp., Sano, Gifu City, Japan.. irig@ghen.co.jp
 SOURCE: GASTROENTEROLOGY, (2000 Aug) 119 (2) 358-67.
 Journal code: FH3; 0374630. ISSN: 0016-5085.
 PUB. COUNTRY: United States
 Journal; Article; (JOURNAL ARTICLE)
 LANGUAGE: English
 FILE SEGMENT: Abridged Index Medicus Journals; Priority Journals
 ENTRY MONTH: 200008
 ENTRY DATE: Entered STN: 20000901
 Last Updated on STN: 20000901
 Entered Medline: 20000822

AB BACKGROUND & AIMS: The significance of acid-primed recognition of ligands by ***Helicobacter*** ***pylori*** ***urease*** is unknown. This study aimed to further characterize the specificity of ***urease*** adherence in vitro and verify whether specific inhibition will translate into in vivo suppression of colonization. METHODS: A highly sensitive competitive enzyme-linked ligand capture assay was used to quantify the capacity of each test inhibitor to compete with labeled mucin for binding sites on immobilized native ***urease***. A model polymer that strongly bound ***urease*** was used in an in vivo trial using euthymic hairless mice as an infection model. RESULTS: The blockage of ***urease***-gastric mucin interaction by certain inhibitors revealed an acid-functional lectin-like activity by ***urease***, specifically recognizing bacterial lipopolysaccharides and certain species of polysaccharides, nonbacterial glycolipids, and ***glycoproteins***. Dextran sulfate significantly ($P < 0.01$) suppressed colonization of mice by H. pylori when given before and/or after challenge. CONCLUSIONS: The acid-driven high-affinity adherence of H. pylori ***urease*** to mucin and lipopolysaccharides contributes to gastric mucosal colonization by the bacterium based on in vivo targeting experiments using specific polysaccharides in a mouse model with acute infection. Acid-functional ***urease***-homing polysaccharides that can interfere with ***urease***-mucin or H. pylori whole cell-mucin interaction in vitro can significantly interfere with colonization by the bacterium in vivo.

L5 ANSWER 3 OF 9 SCISEARCH COPYRIGHT 2002 ISI (R)
 ACCESSION NUMBER: 1999:863987 SCISEARCH
 THE GENUINE ARTICLE: 253AT
 TITLE: Live attenuated Salmonella: a paradigm of mucosal vaccines
 AUTHOR: Sirard J C (Reprint); Niedergang F; Kraehenbuhl J P
 CORPORATE SOURCE: UNIV LAUSANNE, SWISS INST EXPT CANC RES, CH-1066
 EPALINGES, SWITZERLAND (Reprint); UNIV LAUSANNE, INST
 BIOCHEM, CH-1066 EPALINGES, SWITZERLAND
 COUNTRY OF AUTHOR: SWITZERLAND
 SOURCE: IMMUNOLOGICAL REVIEWS, (OCT 1999) Vol. 171, pp. 5-26.
 Publisher: MUNKSGAARD INT PUBL LTD, 35 NORRE SOGADE, PO
 BOX 2148, DK-1016 COPENHAGEN, DENMARK.
 ISSN: 0105-2896.
 DOCUMENT TYPE: General Review; Journal
 FILE SEGMENT: LIFE
 LANGUAGE: English
 REFERENCE COUNT: 211

ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS

AB Two key steps control immune responses in mucosal tissues: the sampling and transepithelial transport of antigens, and their targeting into professional antigen-presenting cells in mucosa-associated lymphoid

tissue. Live *Salmonella* bacteria use strategies that allow them to cross the epithelial barrier of the gut, to survive in antigen-presenting cells where bacterial antigens are processed and presented to the immune cells, and to express adjuvant activity that prevents induction of oral tolerance. Two *Salmonella* serovars have been used as vaccines or vectors, *S. typhimurium* in mice and *S. typhi* in humans. *S. typhimurium* causes gastroenteritis in a broad host range, including humans, while *S. typhi* infection is restricted to humans. Attenuated *S. typhimurium* has been used successfully in mice to induce systemic and mucosal responses against more than 60 heterologous antigens. This review aims to revisit *S. typhimurium*-based vaccination, as an alternative to *S. typhi*, with special emphasis on the molecular pathogenesis of *S. typhimurium* and the host response. We then discuss how such knowledge constitutes the basis for the rational design of novel live mucosal vaccines.

L5 ANSWER 4 OF 9 CAPLUS COPYRIGHT 2002 ACS DUPLICATE 2

ACCESSION NUMBER: 1997:98060 CAPLUS
DOCUMENT NUMBER: 126:198216
TITLE: Inhibition of *Helicobacter pylori* binding to gastrointestinal epithelial cells by sialic acid-containing oligosaccharides
AUTHOR(S): Simon, P. M.; Goode, P. L.; Mobasser, A.; Zopf, D.
CORPORATE SOURCE: Neose Technologies, Inc., Horsham, PA, 19044, USA
SOURCE: Infect. Immun. (1997), 65(2), 750-757
CODEN: INFIBR; ISSN: 0019-9567
PUBLISHER: American Society for Microbiology
DOCUMENT TYPE: Journal
LANGUAGE: English

AB ****Helicobacter**** ****pylori****, the ulcer pathogen residing in the human stomach, binds to epithelial cells of the gastric antrum. The authors have examined binding of 13 bacterial isolates to epithelial cell lines by using a sensitive microtiter plate method in which measurement of bacterial ***urease*** activity provides the means for quantitation of bound organisms. Several established human gastrointestinal carcinoma cell lines grown as monolayers were compared for suitability in these assays, and the duodenum-derived cell line HuTu-80 was selected for testing bacterial binding inhibitors. When bacteria are pretreated with oligosaccharides, ***glycoproteins***, and glycolipids, a complex picture of bacterial-epithelial adherence specificities emerges. Among the monovalent inhibitors tested, 3'-sialyllactose (NeuAc.alpha.2-3Gal.beta.1-4Glc;3'SL) was the most active oligosaccharide, inhibiting adherence for recent clinical isolates of *H. pylori* with a millimolar 50% inhibitory concentration (IC50). Its .alpha.2-6 isomer (6'SL) was less active. Most of the recent clinical isolates examined were inhibited by sialyllactose, whereas long-passaged isolates were insensitive. Among the long-passaged bacterial strains whose binding was not inhibited by 3'SL was the strain ATCC 43504, also known as NCTC 11637 and CCUG 17874, in which the proposed sialyllactose adhesin was recently reported to lack surface expression. Pretreatment of the epithelial monolayer with neuraminidase reduced the extent of binding by those bacteria that are sensitive to inhibition by 3'SL. Other potent inhibitors of bacterial binding are the ***glycoproteins***, .alpha.1-acid ***glycoprotein***, fetuin, porcine gastric and bovine submaxillary mucins, and the glycolipid sulfatide, all of which present multivalent sialylated and/or sulfated galactosyl residues under the conditions of the binding assay. Consistent with this pattern, a multivalent neoglycoconjugate containing 20 mol. of 3'SL per mol. of human serum albumin inhibited bacterial binding with micromolar IC50. The *H. pylori* isolate most sensitive to inhibition by 3'SL was least sensitive to inhibition by sulfatide, gastric mucin, and other sulfated oligosaccharides. Bacteria that have been allowed to bind epithelial cells are also effectively detached by 3'SL. These results describe a heterogeneous adherence repertoire for these bacteria, but they also confirm the critical role of the 3'SL structure on human gastric epithelial cells as an adherence ligand for recent isolates of *H. pylori*.

L5 ANSWER 5 OF 9 MEDLINE DUPLICATE 3

ACCESSION NUMBER: 97256726 MEDLINE
DOCUMENT NUMBER: 97256726 PubMed ID: 9099625
TITLE: Sulfatides inhibit binding of *Helicobacter pylori* to the gastric cancer Kato III cell line.
AUTHOR: Wadstrom T; Hirno S; Novak H; Guzman A; Ringner-Pantzar M; Utt M; Aleljung P

CORPORATE SOURCE: Department of Medical Microbiology, University of Lund,
Solvegatan 23, Lund S-22362, Sweden.
SOURCE: CURRENT MICROBIOLOGY, (1997 May) 34 (5) 267-72.
Journal code: BMJ; 7808448. ISSN: 0343-8651.
PUB. COUNTRY: United States
Journal; Article; (JOURNAL ARTICLE)
LANGUAGE: English
FILE SEGMENT: B
ENTRY MONTH: 199706
ENTRY DATE: Entered STN: 19970612
Last Updated on STN: 19990129
Entered Medline: 19970602

AB ***Helicobacter*** ***pylori*** adhere to Kato III and Hela S3
cells in monolayer cultures. To explore whether cell surface
glycoconjugates on these two cell lines mediate binding of H. pylori,
various carbohydrates, ***glycoproteins***, and glycolipids were
tested to inhibit H.pylori cell adhesion. The adhesion was measured (i)
with a ***urease***-based assay and (ii) by cells stained with
fluorescein. Sodium periodate and sialidase treatment (but not alpha- or
beta-galactosidase, heparitinase, lysozyme, or trypsin) inhibited H. pylori
binding to both cell lines. Sulfatides and sulfated glycoconjugates (50
microg/ml) but not heparin or a number of simple carbohydrates inhibited
binding (1 mg/ml). The two H.pylori strains studied (CCUG 17874 and strain
25) showed high binding of soluble 125I-labeled heparin and other sulfated
carbohydrate compounds.

L5 ANSWER 6 OF 9 EMBASE COPYRIGHT 2002 ELSEVIER SCI. B.V.

ACCESSION NUMBER: 92165602 EMBASE
DOCUMENT NUMBER: 1992165602
TITLE: Antibodies against Helicobacter pylori inhibit the adhesion
of this organism to the gastric mucosal surface.
AUTHOR: Tanaka N.; Kuwayama H.; Sunairi M.; Nakajima M.
CORPORATE SOURCE: Dept. of Internal Medicine (III), School of Medicine, Nihon
University, 1-7-3 Kandasurugadai, Chiyoda-ku, Tokyo 101,
Japan
SOURCE: European Journal of Gastroenterology and Hepatology, (1992)
4/SUPPL. 1 (S67-S69).
ISSN: 0954-691X CODEN: EJGHES
COUNTRY: United Kingdom
DOCUMENT TYPE: Journal; Conference Article
FILE SEGMENT: 004 Microbiology
048 Gastroenterology
LANGUAGE: English
SUMMARY LANGUAGE: English

AB Objectives. The adhesion properties of a strain of ***Helicobacter***
pylori were studied in an in vitro system. Design. H. pylori
adhesion to the gastric mucosal surface was examined in an in vitro
system, using polystyrene assay plates coated with gastric mucus derived
from various sources. The adhesion was further studied using fractions
from mucin, glycosylated beads and anti-H. pylori antibodies. Method. The
numbers of H. pylori cells adhering to the plates were estimated by
measuring ***urease*** activity. Results. There was strong adhesion to
partly purified mucin from a porcine stomach, but only weak adhesion to
mucus derived from cattle. H. pylori adhered to galactosylated beads,
acidic ***glycoprotein*** and the glycolipid components of porcine
mucin. Bacterial adhesion was inhibited not only by whole molecules but
also by the antigen-binding fragment of anti-H. pylori immunoglobulin G.
Conclusions. H. pylori appeared to have an affinity for galactosylated
beads. Further, we suggest that the use of antibodies might be helpful in
treating the H. pylori infection.

L5 ANSWER 7 OF 9 MEDLINE

DUPLICATE 4.

ACCESSION NUMBER: 92003478 MEDLINE
DOCUMENT NUMBER: 92003478 PubMed ID: 1912416
TITLE: Virulence and pathogenicity of Helicobacter pylori.
AUTHOR: Marshall B J
CORPORATE SOURCE: Department of Internal Medicine, University of Virginia
Health Sciences Center, Charlottesville 22908.
SOURCE: JOURNAL OF GASTROENTEROLOGY AND HEPATOLOGY, (1991 Mar-Apr)
6 (2) 121-4. Ref: 28
Journal code: A6J; 8607909. ISSN: 0815-9319.
PUB. COUNTRY: Australia

Journal; Article; (JOURNAL ARTICLE)
General Review; (REVIEW)
(REVIEW, TUTORIAL)

LANGUAGE: English
FILE SEGMENT: Priority Journals
ENTRY MONTH: 199111
ENTRY DATE: Entered STN: 19920124
Last Updated on STN: 19920124
Entered Medline: 19911108

AB H. pylori is a highly virulent organism as evidenced by its low infective dose and widespread high prevalence in human populations. Its virulence is achieved through its ability to survive in a moist environment and its massive ***urease*** production which allows it to survive in the acidic gastric juice long enough to colonize the gastric mucus. Gastric colonization is facilitated by cell wall associated lectins which permit the bacterium to bind to gastric mucus and the gastric epithelial cell. Once in this location, H. pylori produces several enzymes which may harm the gastric epithelium, particularly ***urease*** (through ammonia generation) and phospholipases A and C. H. pylori also weakens the gastric mucous layer by digesting its ***glycoproteins*** and lipids, making the mucus less hydrophobic and more water soluble. ***Helicobacter*** ***pylori*** attracts phagocytic cells, inducing both acute and chronic inflammation as well as an antibody response. Persistence of H. pylori in the mucosa may be enhanced by its cytotoxin and catalase production, by which it survives after phagocytosis by neutrophils.

L5 ANSWER 8 OF 9 MEDLINE DUPLICATE 5

ACCESSION NUMBER: 91147586 MEDLINE
DOCUMENT NUMBER: 91147586 PubMed ID: 1997534
TITLE: Breakdown of gastric mucus in presence of Helicobacter pylori.
AUTHOR: Sidebotham R L; Batten J J; Karim Q N; Spencer J; Baron J H
CORPORATE SOURCE: Department of Surgery, Royal Postgraduate Medical School, Hammersmith Hospital, London.
SOURCE: JOURNAL OF CLINICAL PATHOLOGY, (1991 Jan) 44 (1) 52-7.
Journal code: HT3; 0376601. ISSN: 0021-9746.
PUB. COUNTRY: ENGLAND: United Kingdom
Journal; Article; (JOURNAL ARTICLE)
LANGUAGE: English
FILE SEGMENT: Abridged Index Medicus Journals; Priority Journals
ENTRY MONTH: 199104
ENTRY DATE: Entered STN: 19910419
Last Updated on STN: 19910419
Entered Medline: 19910404

AB The potential of ***Helicobacter*** ***pylori*** to degrade gastric mucus was examined. Colonies of H pylori cultured from antral mucosal biopsy specimens of patients with non-autoimmune gastritis were washed with sterile saline, passed through a sterilisation filter, and the filtrate examined for ***urease***, protease, and mucolytic activity. The filtrate failed to hydrolyse bovine serum albumin, or to degrade stable mucus ***glycoprotein*** structures of high particle weight that had been separated from human gastric mucus on Sepharose 2B. The high particle weight mucus ***glycoprotein*** was, however, extensively degraded when incubated with H pylori filtrate (which possessed ***urease*** activity) in the presence of 2 M urea, to release fragments of Mr approximately 2×10^6 . The high particle weight mucus ***glycoprotein*** was also broken down to a comparable extent when incubated with Jack bean ***urease*** in the presence of 2 M urea, or 1 M ammonium carbonate, or 40 mM carbonate-bicarbonate buffer (pH 8.7), but not when treated with 4 M urea alone, or Jack bean ***urease*** alone. These results indicate that the loss of high particle weight mucus ***glycoprotein*** in gastric mucus from patients with gastritis and gastric ulcers is unlikely to be due to the mucolytic action of an extra-cellular protease produced by H pylori, but it may result from the destabilising effects of a carbonate-bicarbonate buffer, generated at the mucosal surface when H pylori ***urease*** hydrolyses transuded plasma urea.

L5 ANSWER 9 OF 9 MEDLINE
ACCESSION NUMBER: 90306646 MEDLINE
DOCUMENT NUMBER: 90306646 PubMed ID: 2194877
TITLE: [Does Helicobacter pylori have a direct proteolytic effect

in ulcerative disease?].
L'Helicobacter pylori esercita un'azione proteolitica
diretta in corso di malattia ulcerosa?.

AUTHOR: Tessaro P; Di Mario F; Vianello F; Dal Santo P; Germana B;
Plebani M; Faggian D; Del Favero G; Naccarato R
CORPORATE SOURCE: Cattedra e Divisione di Gastroenterologia, Università di
Padova.
SOURCE: GIORNALE DI CLINICA MEDICA, (1990 Mar) 71 (3) 173-8, 181.
Journal code: FAP; 0413411. ISSN: 0017-0275.
PUB. COUNTRY: Italy
Journal; Article; (JOURNAL ARTICLE)
LANGUAGE: Italian
FILE SEGMENT: Priority Journals
ENTRY MONTH: 199008
ENTRY DATE: Entered STN: 19900921
Last Updated on STN: 19900921
Entered Medline: 19900816

AB ***Helicobacter*** ***pylori*** (H.p.), has been shown,
experimentally, to exert a proteolytic activity against mucous fractions.
Aim of this study was to assess the prevalence of H.p. in peptic ulcer and
to analyze its possible influence on gastric mucus components, on peptic
activity in gastric juice and the possible action on peptic secretion. 223
patients undergoing upper gastrointestinal endoscopy were analyzed for the
presence of H.p. in the mucosa: 99 duodenal ulcer patients (D.U.), 58
gastric ulcer patients (D.U.) and 66 dyspeptic subjects. In each patients,
three contiguous gastric biopsies were taken at the antrum: the first was
evaluated for gastritis (Whitehead Criteria), the two other analyzed for
H.p. with a rapid ***urease*** test. In a subgroup of 25 D.U. and 18
G.U. patients, two other biopsies were taken at the fundus corpus of the
stomach, to evaluate peptic secretion. To determinate mucous components
(acid and neutral ***glycoproteins***, galactose and
N-acetylneuraminic acid), gastric juice samples were collected during
endoscopy. H.p. was present in 89% of antral biopsies in D.U., in 56% of
G.U. and in 51% of D., and was associated to antral gastritis. As regard
gastric juice components, we observed an increase and a decrease of acid
glycoproteins, respectively; in D.U. and G.U. patients with H.p.
infection. An increase of peptic activity has been found in the gastric
juice of both gastric and duodenal ulcer patients H.p. positive (G.U. p
less than 0.05). On the contrary, no significant differences were observed
on peptic activity in the fundus-corpus biopsies between H.p. positive and
H.p. negative patients. (ABSTRACT TRUNCATED AT 250 WORDS)

=> d his

(FILE 'HOME' ENTERED AT 15:21:03 ON 24 MAR 2002)

FILE 'MEDLINE, CAPLUS, BIOSIS, EMBASE, SCISEARCH, AGRICOLA' ENTERED AT
15:21:34 ON 24 MAR 2002

L1 497584 S GLYCOPROTEIN
L2 77272 S HELICOBACTER PYLORI
L3 6694 S L2 (P) UREASE
L4 24 S L1 (P) L3
L5 9 DUPLICATE REMOVE L4 (15 DUPLICATES REMOVED)

=> s l2 (p) colonization (p) inhibit?
L6 403 L2 (P) COLONIZATION (P) INHIBIT?

=> s l6 and l1
L7 20 L6 AND L1

=> duplicate remove l7
DUPLICATE PREFERENCE IS 'MEDLINE, CAPLUS, BIOSIS, EMBASE, SCISEARCH'
KEEP DUPLICATES FROM MORE THAN ONE FILE? Y/(N):n
PROCESSING COMPLETED FOR L7
L8 8 DUPLICATE REMOVE L7 (12 DUPLICATES REMOVED)

=> s l8 not l5
L9 6 L8 NOT L5

=> d l9 1-6 ibib abs

L9 ANSWER 1 OF 6 MEDLINE
 ACCESSION NUMBER: 2000158106 MEDLINE
 DOCUMENT NUMBER: 20158106 PubMed ID: 10695559
 TITLE: Helicobacter pylori lipopolysaccharide-mediated gastric and extragastric pathology.
 AUTHOR: Moran A P
 CORPORATE SOURCE: Department of Microbiology, National University of Ireland. Galway.. anthony.moran@nuigalway.ie
 SOURCE: JOURNAL OF PHYSIOLOGY AND PHARMACOLOGY, (1999 Dec) 50 (5) 787-805. Ref: 97
 Journal code: A9B; 9114501. ISSN: 0867-5910.
 PUB. COUNTRY: Poland
 Journal; Article; (JOURNAL ARTICLE)
 General Review; (REVIEW)
 (REVIEW, TUTORIAL)
 LANGUAGE: English
 FILE SEGMENT: Priority Journals
 ENTRY MONTH: 200003
 ENTRY DATE: Entered STN: 20000407
 Last Updated on STN: 20000407
 Entered Medline: 20000328

AB Lipopolysaccharides (LPS) are a family of toxic phosphorylated glycolipids in the outer membrane of Gram-negative bacteria, including
 Helicobacter ***pylori***, and are composed of a lipid moiety (termed lipid A), a core oligosaccharide, and a polymeric O-specific polysaccharide chain. Compared with LPS of other bacteria, H. pylori LPS and lipid A induce low immunological activities in a range of test systems. Nevertheless, these reduced levels of LPS-induced cytokines and toxic oxygen radicals can contribute, with those induced by bacterial proteins, to the H. pylori-associated inflammatory response. Whether the ability of H. pylori LPS to induce low production of both procoagulant activity and plasminogen activator ***inhibitor*** type 2 by human mononuclear cells contributes to localized inflammatory responses alone and, in addition, play a role in extragastric pathology remains an open question. The core oligosaccharide of H. pylori LPS, in part with a 25 kDa protein adhesin, mediates the binding of the bacterium to the host ***glycoprotein*** laminin, and hence interferes with gastric cell receptor-laminin interaction in the basement membrane. Also affecting mucosal integrity, the core sugars of certain H. pylori strains, particularly those associated with gastric ulceration, have been implicated in pepsinogen induction, but this is a strain-dependent phenomenon. Of particular interest, the O-chains of a large proportion of H. pylori strains mimic Lewis (Le) antigens. Although investigations have focussed on the role of these antigens in H. pylori-associated autoimmunity, which remains to be unequivocally established, other pathogenic consequences of Lewis mimicry are becoming apparent. Expression of Lewis antigens may be crucial for H. pylori ***colonization*** and adherence and, by aiding bacterial interaction with the gastric mucosa, thereby aid delivery of secreted products, and hence influence the inflammatory response.

L9 ANSWER 2 OF 6 MEDLINE
 ACCESSION NUMBER: 92391434 MEDLINE
 DOCUMENT NUMBER: 92391434 PubMed ID: 1381553
 TITLE: Glycosulfatase activity of H. pylori toward human gastric mucin: effect of sucralfate.
 AUTHOR: Slomiany B L; Murty V L; Piotrowski J; Grabska M; Slomiany A
 CORPORATE SOURCE: Research Center, New Jersey Dental School, University of Medicine and Dentistry of New Jersey, Newark.
 CONTRACT NUMBER: AA05858-11 (NIAAA)
 DK31684-15 (NIDDK)
 SOURCE: AMERICAN JOURNAL OF GASTROENTEROLOGY, (1992 Sep) 87 (9) 1132-7.
 Journal code: 3HE; 0421030. ISSN: 0002-9270.
 PUB. COUNTRY: United States
 Journal; Article; (JOURNAL ARTICLE)
 LANGUAGE: English
 FILE SEGMENT: Priority Journals
 ENTRY MONTH: 199210
 ENTRY DATE: Entered STN: 19921023
 Last Updated on STN: 19960129

AB ***Colonization*** of gastric mucosa by ***Helicobacter pylori***, a bacterium implicated in the etiology of gastric disease, involves the cell surface sulfated glycosphingolipid receptors for the attachment. Evidence has also been obtained recently that sulfated mucus ***glycoproteins*** have the ability to interfere with this process. Here, we show that *H. pylori* displays glycosulfatase activity, and report the specificity of this enzyme toward gastric mucosal sulfated ***glycoproteins*** and glycolipids. With 35S-labeled human gastric sulfated mucin as substrate, the enzyme activity was identified in the extracellular material elaborated by the bacterium. The glycosulfatase exhibited maximum activity at pH 5.7 in the presence of Triton X-100 and CaCl₂, and gave on SDS-PAGE a protein band of 30 kDa. Specificity studies revealed that the enzyme effectively caused desulfation of N-acetylglucosamine-6-sulfate and galactose-6-sulfate present in carbohydrate chains of gastric mucins, as well as that of glucose-6-sulfate, a constituent of mucus glyceroglucolipids. However, the *H. pylori* glycosulfatase was ineffective toward galactosyl- and lactosylceramide sulfates which serve as receptors for this bacterium attachment and contain the sulfate ester group at C-3 of galactose. The glycosulfatase activity toward human sulfated gastric mucin was ***inhibited*** by sucralfate. The ***inhibitory*** effect was proportional to the concentration of sucralfate up to 120 micrograms/ml, at which a 78% decrease in mucin desulfation occurred. The results demonstrate that *H. pylori*, through its glycosulfatase activity, affects the sulfated mucin and glyceroglucolipid content of the protective mucus layer, and that antiulcer drug sucralfate is able to counteract the detrimental action of this enzyme.

L9 ANSWER 3 OF 6 EMBASE COPYRIGHT 2002 ELSEVIER SCI. B.V.
 ACCESSION NUMBER: 2001192176 EMBASE
 TITLE: Safe as mother's milk: Carbohydrates as future anti-adhesion drugs for bacterial diseases.
 AUTHOR: Sharon N.; Ofek I.
 CORPORATE SOURCE: N. Sharon, Department of Biological Chemistry, Weizmann Institute of Science, Rehovot 76100, Israel.
 bfsharon@weizmann.weizmann.ac.il
 SOURCE: Glycoconjugate Journal, (2000) 17/7-9 (659-664).
 Refs: 24
 ISSN: 0282-0080 CODEN: GLJOEW
 COUNTRY: Netherlands
 DOCUMENT TYPE: Journal; General Review
 FILE SEGMENT: 030 Pharmacology
 037 Drug Literature Index
 LANGUAGE: English
 SUMMARY LANGUAGE: English

AB The majority of infectious diseases are initiated by adhesion of pathogenic organisms to the tissues of the host. In many cases, this adhesion is mediated by lectins present on the surface of the infectious organism that bind to complementary carbohydrates on the surface of the host tissues. Lectin-deficient mutants often lack ability to initiate infection. Soluble carbohydrates recognized by the bacterial lectins block the adhesion of the bacteria to animal cells in vitro. Moreover, they have also been shown to protect against experimental infection by lectin-carrying bacteria in different organs of mammals such as mice, rabbits, calves and monkeys. In a phase II clinical trial, a pentasaccharide shown to have anti-adhesive activity against *Streptococcus pneumoniae* and *Haemophilus influenzae* in vitro failed to protect young children from nasopharyngeal ***colonization*** with these organisms and from developing otitis media. This could be because insufficient drug was delivered via nasal spray, because bacteria express multiple specificities, the ***inhibition*** of which may require a cocktail of oligosaccharides, or because children have different carbohydrate receptors from those of adults. The results of a clinical trial in which N-acetylneuraminyl(.alpha.2-3)lactose was administered orally to ***Helicobacter pylori*** positive patients in an effort to reduce or eradicate bacterial ***colonization***, are awaited with interest. Although the high cost of production of the required oligosaccharides is falling with the recent introduction of enzymatic methods of synthesis, new technologies, in particular the use of engineered bacteria, promise to lower it even further. Attachment of the oligosaccharides to soluble polymeric carriers will increase greatly their

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ACCESSION NUMBER: 93:283735 SCISEARCH

THE GENUINE ARTICLE: KZ690

TITLE: ***INHIBITION*** OF ***HELICOBACTER*** -
PYLORI ***COLONIZATION*** BY AN ANTIULCER
AGENT, SULGLYCOTIDE
AUTHOR: CZAJKOWSKI A (Reprint); PIOTROWSKI J; YOTSUMOTO F;
SLOMIANY A; SLOMIANY B L
CORPORATE SOURCE: UNIV MED & DENT NEW JERSEY, RES CTR, NEWARK, NJ, 07103
(Reprint)
COUNTRY OF AUTHOR: USA
SOURCE: BIOCHEMISTRY AND MOLECULAR BIOLOGY INTERNATIONAL, (APR
1993) Vol. 29, No. 5, pp. 965-971.
ISSN: 1039-9712.
DOCUMENT TYPE: Article; Journal
FILE SEGMENT: LIFE
LANGUAGE: ENGLISH
REFERENCE COUNT: 27

L9 ANSWER 6 OF 6 SCISEARCH COPYRIGHT 2002 ISI (R)

ACCESSION NUMBER: 91:596194 SCISEARCH

THE GENUINE ARTICLE: GL944

TITLE: ***INHIBITION*** OF ***HELICOBACTER*** -
PYLORI ***COLONIZATION*** BY SULFATED GASTRIC
MUCIN
AUTHOR: PIOTROWSKI J (Reprint); SLOMIANY A; MURTY V L N; FEKETE Z;
SLOMIANY B L
CORPORATE SOURCE: UNIV MED & DENT NEW JERSEY, NEW JERSEY DENT SCH, RES CTR,
NEWARK, NJ, 07103 (Reprint)
COUNTRY OF AUTHOR: USA
SOURCE: BIOCHEMISTRY INTERNATIONAL, (1991) Vol. 24, No. 4, pp.
749-756.
DOCUMENT TYPE: Article; Journal
FILE SEGMENT: LIFE
LANGUAGE: ENGLISH
REFERENCE COUNT: 26

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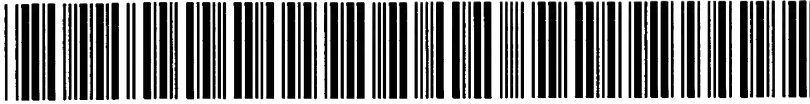
L1 497584 S GLYCOPROTEIN
L2 77272 S HELICOBACTER PYLORI
L3 6694 S L2 (P) UREASE
L4 24 S L1 (P) L3
L5 9 DUPLICATE REMOVE L4 (15 DUPLICATES REMOVED)
L6 403 S L2 (P) COLONIZATION (P) INHIBIT?
L7 20 S L6 AND L1
L8 8 DUPLICATE REMOVE L7 (12 DUPLICATES REMOVED)
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